

PD-L1 [Biotinylated]: PD-1 Inhibitor Screening ELISA Kit

Pack Size: 96 tests

Catalog Number: EP-158

IMPORTANT: Please carefully read this manual before performing your experiment.

For Research Use Only. Not For Use In Diagnostic Or Therapeutic Procedures

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INTENDED USE

This kit is designed for screening of inhibitors of binding between human PD-L1 and human PD-1.

It is intended for research use only (RUO).

PRINCIPLE OF THE ASSAY

This inhibitor screening ELISA kit is designed to facilitate the identification and characterization of new PD-L1

pathway inhibitors. The assay takes advantage of our in house-developed binding of biotinylated human PD-L1 to

immobilized human PD-1 in a functional ELISA assay and employs a simple colorimetric ELISA platform. Briefly,

we provide you with a human Biotinylated PD-L1 protein, a human PD-1 protein, an anti-PD-L1 neutralizing

antibody (as method verified Std.), and Streptavidin-HRP reagent. Your experiment will include 4 simple steps:

1) Coat the plate with human PD-1.

2) Add your molecule of interest to the tests.

3) Add human PD-L1-Biotin to bind the coated human PD-1.

4) Add Streptavidin-HRP followed by TMB or other colorimetric HRP substrate.

Finally, the half maximal inhibitory concentration (IC50) of your compound to PD-L1: PD-1 binding will be

determined by comparing OD readings among different experimental groups.

MATERIALS PROVIDED

TABLE 1. MATERIALS PROVIDED (pls modify according to COA)

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Catalog	Components	Size (96 tests)	Format	Storage	
EP158-C01	High-bind Plate	1 plate	Solid	2-8℃	
EP158-C02	Human PD-1	50 μg	Powder	2-8°C	
EP158-C03	Biotinylated Human PD-L1	10 µg	Powder	2-8°C	-70°C after
EP158-C04	Anti-PD-L1 Neutralizing Antibody	20 μg	Powder	2-8°C	reconstitution, avoid freeze-thaw cycles
EP158-C05	Streptavidin-HRP	5 μg	Powder	2-8°C, avoid light	110020 111111 0,000
EP158-C06	Coating Buffer	12 mL	Liquid	2-8°C	
EP158-C07	10xWashing Buffer	50 mL	Liquid	2-8°C	
EP158-C08	Blocking Buffer	50 mL	Liquid	2-8°C	
EP158-C09	Substrate Solution	12 mL	Liquid	2-8°C, avoid light	
EP158-C10	Stop Solution	7 mL	Liquid	2-8°C	

REAGENTS/EQUIPMENT NEEDED BUT NOT SUPPLIED

Single or dual wavelength microplate reader with 450 nm/630nm filter;

Centrifuge;

37 °C Incubator;

Single channel or multichannel pipettes with 10 μ L, 200 μ L and 1000 μ L precision;

 $10 \mu L$, $200 \mu L$ and $1000 \mu L$ pipette tips;

Test Tubes:

Graduated cylinder;

Deionized or distilled water for dilution;

STORAGE AND VALIDITY INSTRUCTIONS

Unopened kit should be stored at 2°C -8°C upon receiving. Find the expiration date on the outside packaging and do not use reagents past their expiration date.

The kit should be stored as TABLE 1 after the reconstitution of lyophilized materials. The shelf life is 30 days from

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the date of opening.

Note:

- a. Do not use reagents past their expiration date.
- b. Find the expiration date on the outside packaging.

REAGENT PREPARATION

- 1. Restore all reagents and samples to room temperature (20-25°C) before use.
- 2. Reconstitute the provided lyophilized materials to stock solutions with sterile deionized water as recommended in Tab.2, Solubilize for 15 to 30 minutes at room temperature with occasional gentle mixing. Avoid vigorous shaking or vortex. The reconstituted stock solutions should be stored at -70°C. **Avoid freeze-thaw cycles**.

Note: Streptavidin-HRP stock solution should be protected from light.

TABLE 2. RECONSTITUTION METHODS FOR 96 TESTS

Catalog	Components	Amount	Stock Solution Con.	Reconstitution Buffer and Vol.
EP158-C02	Human PD-1	50 μg	100 μg/mL	500 μL, water
EP158-C03	Biotinylated Human PD-L1	10 μg	100 μg/mL	100 μL, water
EP158-C04	Anti-PD-L1 Neutralizing Antibody	20 μg	100 μg/mL	200 μL, water
EP158-C05	Streptavidin-HRP	5 μg	50 μg/mL	100 μL, water

RECOMMENDED PROTOCOL

1. Working solution preparation

1.1 Preparation of 1×Washing Buffer:

Dilute 50 mL 10×Washing Buffer with ultrapure water/deionized water to 500 mL.

1.2 Preparation of Dilution Buffer:

10 mL Blocking Buffer (EP158-C08) add 30 mL 1×Washing Buffer.

2. Coating

1)Dilute Human PD-1 stock solution (100 μg/mL) to 4 μg/mL with Coating Buffer to make Human PD-1 working

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solution.

2)Add 100 µL of Human PD-1 working solution (4 µg/mL) to each well and leave a couple of wells uncoated for No-

Coating Control, seal the plate with microplate sealing film and incubate overnight (or 16 hours) at 4°C.

3. Washing

Remove the remaining solution by aspiration, add 300 μL of 1×Washing Buffer to each well, gently tap the plate for

1 minute, remove any remaining 1×Washing Buffer by aspirating or decanting, invert the plate and blot it against

paper towels. Repeat the washing step above for three times.

Note: For best results, the complete removal of the Human PD-1 solution is essential. The use of a manifold dispenser or an auto-washer

may be necessary.

4. Blocking

Add 300 µL Blocking Buffer to each well, seal the plate with microplate sealing film and incubate at 37°C for 1.5

hours.

5. Washing

Repeat step 3. At the same time, you can start to prepare your samples.

6. Add Samples

1)Make serial dilution of the samples as appropriate.

2)If you intend to use the provided Anti-PD-L1 Neutralizing Antibody as a reference (Std.), you may dilute the

antibody as recommended in Figure 1.

3)Add 50 µL of sample solution to each well according to our recommendation (Figure 2) or your own plate setup.

4)For No-Coating Control wells, please add 50 µL Dilution Buffer.

7.Binding

1) Dilute Biotinylated Human PD-L1 stock solution (100 µg/mL) to 0.06 µg/mL with Dilution Buffer to make

Biotinylated Human PD-L1 working solution.

2) For No-binding control wells, please add 50 µL Dilution Buffer.

3) For all other wells, please add 50 µL Biotinylated Human PD-L1 working solution to the wells and mix the samples

by gently tapping the plate. Seal the plate with microplate sealing film and incubate at 37°C for 1 hour.

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Note: The working solution should be prepared immediately before use and should not be stored.

FIG.1 PREPARATION OF 1:1 SERIAL DILUTIONS OF THE Anti-PD-L1 Neutralizing Antibody

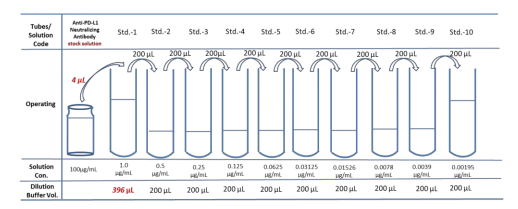


FIG.2 PLATE LAYOUT

	1 2	2 3	4	5	6	7	8	9	10	11	12
А	Std8 St	d8 Std9	Std9)())())	
В	Std7 St	d7 Std10	Std10)()()()()()()	
С	Std6 St	d6 Positive Ctrl.	Positive Ctrl.)()()()()()()	
D	Std5 St	d5 No- binding Ctrl.	No- binding Ctrl.)()()()()()()	
E	Std4 St	d4 No- coating Ctrl.	No- coating Ctrl.)()()()()()()	
F	Std3 St	d3	())()()()()()()	
G	Std2 St	d2	())()()()()()()	
н	Std1	:d1	())()()()()())	

8. Washing

Repeat step 3.

9.Add Streptavidin-HRP

- 1)Dilute Streptavidin-HRP stock solution (50 μg/mL) to 0.1 μg/mL with Dilution Buffer to make Streptavidin-HRP working solution.
- 2)For all wells, add $100 \,\mu\text{L}$ Streptavidin-HRP working solution, seal the plate with microplate sealing film and incubate at 37°C for 1 hour, avoid light.

10.Washing

Repeat step 3.

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11. Substrate Reaction

Add 100 µL **Substrate Solution** to each well. Seal the plate with microplate sealing film and incubate at 37°C for 20 minutes. Avoid light.

12.Termination

Add 50 µL Stop Solution to each well, and gently shake the plate to allow thorough mixing.

Note: the color in the wells should change from blue to yellow.

13.Data Recording

Read the absorbance at 450 nm using UV/Vis microplate spectrophotometer.

Note: Subtracting the value read at OD_{450nm} with OD_{630nm} can be used to reduce the background noise.

SIMPLIFIED PROTOCOL

TABLE. 3 ASSAY PROTOCOL

				F			_
Steps Code	Steps	Reagents & Instruments	Reaction Conditions	Samples	No-binding	No-coating	Positive
					Ctrl.	Ctrl.	Ctrl.
1	Working fluid preparation	N/A	N/A	N/A	N/A	N/A	N/A
2	Coating	Human PD-1 Working Solution	4°C for overnight	100 μL	100 μL	_	100 μL
3	Washing	1XWash Buffer	Wash for 3 times	300 μL	300 μL	300 μL	300 μL
4	Blocking	Blocking Buffer	37°C for 1.5 hours	300 μL	300 μL	300 μL	300 μL
5	Washing	1XWash Buffer	Wash for 3 times	300 μL	300 μL	300 μL	300 μL
6	Add Samples	Samples	_	50 μL	_	_	_
0		Dilution Buffer		_	50 μL	50 μL	50 μL
7	Binding	Biotinylated Human PD-L1 Working Solution	Mix by gentle tapping, incubate at 37°C for 1 hours	50 μL	_	50 μL	50 μL
		Dilution Buffer		_	50 μL	_	_
8	Washing	1XWash Buffer	Wash for 3 times	300 μL	300 μL	300 μL	300 μL
9	Streptavidin-HRP	Streptavidin-HRP Working Solution	37°C for 1 hours	100 μL	100 μL	100 μL	100 μL

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10	Washing	1XWash Buffer	Wash for 3 times	300 μL	300 μL	300 μL	300 μL
11	Substrate Reaction	Substrate Solution	37°C for 20 minutes	100 μL	100 μL	100 μL	100 μL
12	Termination	Stop Solution	Mix by gentle tapping	50 μL	50 μL	50 μL	50 μL
13	Data Recording	UV/Vis spectrophotometer	Measure absorbance at 450 nm, with the correction wavelength set at 630 nm				

Note for TAB. 3:

- 1) Samples: Your samples of interest.
- 2) No-binding Ctrl.: Reaction without Biotinylated Human PD-L1 added. The absorbance should be around 0.05(< 0.1) at 450 nm.
- 3) No-coating Ctrl.: Reaction without Human PD-1 coated on the wells. The absorbance should be around 0.05(< 0.1) at 450 nm.
- 4) Positive Ctrl.: Determined the max value in 450nm absorbance, when out of inhibitors.
- 5) It is recommended that all samples, controls and standards should be done in duplicates.

PRECAUSIONS

- 1. This kit is for research use only and is not for use in diagnostic or therapeutic applications.
- 2. This kit should be used according to the provided instructions.
- 3. Do not mix reagents from different lots.
- 4. All reagents should be balanced to room temperature (20°C-25°C) before use.
- 5. This kit should be stored at 2°C-8°C.
- 6. Please prepare the working solution of each component according to the needs of the experiment. Except for 1x Washing Buffer, all prepared working solution is for one-time use and cannot be stored.

METHOD VERIFICATION

INHIBITION OF PD-L1 [Biotinylated]: PD-1 BINDING BY ANTI-PD-L1 NEUTRALIZING ANTIBODY

Serial dilutions of Anti-PD-L1 Neutralizing antibody (Catalog # EP158-C04) (1:1 serial dilution, from 1 μg/mL to 0.00195μg/mL) was added into PD-L1 [Biotinylated]: PD-1 binding reactions. The assay was performed according to the protocol described below. Background was subtracted from data points prior to log transformation and curve fitting.

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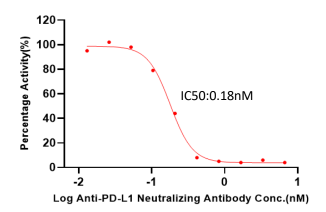
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Anti-PD-L1 Neutralizing Antibody Conc.(µg/ml)	Anti-PD-L1 Neutralizing Antibody Conc.(nM)	Mean Abs.(OD450)	Percentage Activity(%)		
0	0.000	2.469	100%		
0.002	0.013	2.355	95%		
0.004	0.026	2.511	102%		
0.008	0.052	2.425	98%		
0.016	0.104	1.946	79%		
0.031	0.208	1.075	44%		
0.063	0.417	0.202	8%		
0.125	0.833	0.113	5%		
0.250	1.667	0.098	4%		
0.50	3.333	0.142	6%		
1.0	6.667	0.107	4%		
No Coating		0.096			
No Binding		0.071			

For each experiment, a standard curve needs to be set for each micro-plate, and the specific OD value may vary depending on different laboratories, testers, or equipments. The example data is for reference only.

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